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# Pharmaceutical Dosage Forms and Drug Delivery Systems

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Fig. 3-2. As the transport maximum of proposeds a range makesake a representation course on the manda me Arm C Rollb. W.L. Mart. J. Propos. 4.7 ses. Phys. s.

transport as a subclassification of specialized transport, denotes a process with the additional teature of the solute or drug being moved across the membrane against a concentration gradient, that is, from a solution of lower concentration to one of a higher concentration or, if the solute is an ion, against an electrochemical potential gradient. In contrast to active transport, facilitated diffusion is a specialized transport mechanism having all of the above characteristics except that the solute is not transferred against a concentration gradient and may attain the same concentration inside the cell as that on the outside

Many body nutrients, as sugars and amino icids, are transported across the membranes of the gastrointestinal tract by carrier processes. Certain vitamins, as thiamine, niacin, riboflavin and vitamin B<sub>0</sub>, and drug substances as methyl dopa and 5-fluorouracil, require active transport mechanisms for their absorption.

Investigations of intestinal transport have often utilized of sitic (at the site) or in the interpretation of the body) animal models of the control of transport models, however, recently cell culture models of human small-intestine absorptive cells have become available to investigate transport across intestinal epithelium. Both passive and transport-mediated studies have been conducted to investigate mechanisms as well as tates of transport

### Dissolution and Drug Absorption

In order for a drug to be absorbed, it must first be dissolved in the third at the absorption site.

tor instance a drug administered orange as table of capsule form cannot be absorbed until the drug particles are dissolved by the flinds at some point, within the gastrointestinal tract. In instances in which the solubility of a drug is dependent upon either an acidic or basic medium, the drug would be dissolved in the stomach or intestines respectively. (Fig. 3–3). The process by which a drug particle dissolves is termed dissolved treat.

As a drug particle undergoes dissolution, the drug molecules on the surface are the first to enter into solution creating a saturated layer of drug solution which envelops the surface of the solid drug particle. This layer of solution is referred to as the *dimesion layer*. From this diffusion layer, the drug molecules pass throughout the dissolving fluid and make contact with the biologic membranes and absorption ensues. As the molecules of drug continue to leave the diffusion layer, the layer is replenished with dissolved drug from the surface of the drug particle and the process of absorption continues.

If the process of dissolution for a given drug particle is rapid, or if the drug is administered as a solution and remains present in the body as such, the rate at which the drug becomes absorbed would be primarily dependent upon its ability to traverse the membrane barrier. However, if the rate of dissolution for a drug particle



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can a terrary for the string substance or the dosage tories for the diring substance or the dosage tories for the dissolution process useff would be a rate line to gister in the absorption process slowly solution at appears in diagram may not seek be absorbed at a slow rate, they may be up-ompletely absorbed or in some cases targety mabsorbed following or al administration, due to the natural limitation of time that they may remain within the domach or the intestinal tract. Thus, poorly soluble drugs or poorly formulated drug products may result in a drug's incomplete absorption, and its passage, unchanged, out of the system via the teres.

Under normal circumstances a drug may be expected to remain in the stomach for 2 to 4 hours our transportance from and in the small intestines for 4 to 10 hours, although there is substantial variation between people, and even in the same person on different occasions. Various techniques have been used to determine gastric emptying time and the gastrointestinal passage of drug from Various oral dosage forms, includand the tracking of dosage forms labeled with camma emitting fadionuclides through gamma The gastric emptying time for a seintiglaphy. dring is most rapid with a fasting stomach, becoming slower as the food content is increased changes in gastric emptying time and or in inrestinal motility can affect drug transit time and thus the opportunity for drug dissolution and absorption

These changes can be effected by drugs the patient may be taking. Certain drugs with anticholmergic properties, e.g., dicyclomine HCL amitripty line HCL have the ability to slow down gastric emptying. This can enhance the rate of absorption of drugs normally absorbed from the stomach, and reduce the rate of absorption of drugs that are primarily absorbed from the small intestine. Alternatively drugs which enhance gastric motiaty leig. Jaxatives, may cause some dings to move so quickly through the gastroin. testinal system and past their absorptive site at such a rate to reduce the amount of drug actually absorbed. This effect has been demonstrated with digoxin, whose absorption is significantly idecreased by accelerating gastromtestinal moalay

The aging process itself may also influence gastrointestical absorption in the elderly gastra acidity the number of absorptive cells intestinal blood flows the rate of gastra emptying and in restinal motility are all decreased. It appears

or in the forest of which insorption in dependent upon parente processes in not affected by the selfictors is unable. Those that its depention from the firms both mechanisms of good from from the armine steps. A decrease in gasthe emptying time would be advantageous for those drugs that are absorbed from the stomach but disady integeous for those drugs which are prone to acid degradation of gopenicallins, crythtomychic of inactivated by stomach enzymes of dopa.

The dissolution of a substance may be described by the modified Noves Whitney equation

in which do dris the rate of dissolution, k is the dissolution rate constant. S is the surface area of the dissolving solid, cois the saturation concentration of drug in the diffusion layer (which may be approximated by the maximum solubility of the drug in the solvent since the diffusion layer is considered saturated), and d, is the concentration of the drug in the dissolution medium at cois the concentration gradient). The rate of dissolution is governed by the rate of diftusion of solute molecules through the diffusion Liver into the body of the solution. The equation reveals that the dissolution rate of a drug may be increased by increasing the surface area treducing the particle size) of the drug, by increasing the solubility of the drug in the diffusion layer, and by factors embodied in the dissolution. rate constant, k. including the intensity of agitation of the solvent and the diffusion coefficient of the dissolving drug. For a given drug, the diftusion coefficient and usually the concentration or the drug in the diffusion layer will increase with increasing temperature. Also, increasing the rate of aggration of the dissolving medium. will increase the rate of dissolution. A reduction in the viscosity of the solvent employed is another means which may be used to enhance the dissolution rate of a drug. Changes in the pH or the nature of the solvent which influence the solubility of the drug may be used to advantage in increasing dissolution rate. Effervescent, butteach aspirin tablet formulations use some of these principles to their advantage. Due to the aikaline adjuvants in the tablet, the solubility of the ispining enhanced within the diffusional

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